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Results of Search in 1976 to present db for:

AN/pharmacia: 1318 patents.

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Refine Search

an/pharmacia and eplerenone

PAT. NO.	Title
1 6,903,127	T Method of treating sexual disturbances
2 6,903,097	T Heterocycle carboxamides as antiviral agents
3 6,902,889	T Analytical method and advice
4 6,900,244	T Anilino liver X-receptor modulators
5 6,900,232	T Cycloalkyl alkanoic acids as integrin receptor antagonists
6 6,897,318	T Process for making substituted pyrazoles
7 6,894,042	T Azabicyclic compounds for the treatment of disease
8 6,891,025	T Modified cytokine receptor protein
9 6,890,945	T Method of treating sexual disturbances
10 6,890,937	T Aromatic sulfone hydroxamic acid metalloprotease inhibitor
11 6,890,928	T Aromatic sulfone hydroxamic acids and their use as protease inhibitors
12 6,890,920	T Quaternary ammonium compounds
13 6,887,995	T Process to prepare oxazolidinones
14 6,887,868	T Therapeutic 5-HT ligand compounds
15 6,884,813	T Antimicrobial dihydrothiazine and dihydrothiopyran oxazolidinones
16 6,881,833	T Component in the hedgehog signalling pathway
17 6,878,823	T 1,2,3,4,5,6-hexahydroazepino[4,5-b]indoles containing arylsulfones at the 9-position
18 6,878,730	T Quaternary ammonium compounds
19 6,878,516	T Autoregulatory system for validating microbial genes as possible antimicrobial targets using a tetracycline-controllable element
20 6,875,871	T Oxazolidinone photoaffinity probes

- 21 [6,875,791](#) [T](#) [Polycyclic aryl and heteroaryl substituted 4-pyrones useful for selective inhibition of the coagulation cascade](#)
- 22 [6,875,790](#) [T](#) [Cyclic sulfone containing retroviral protease inhibitors](#)
- 23 [6,875,785](#) [T](#) [Heterocyclo substituted hydroxamic acid derivatives as cyclooxygenase-2 and 5-lipoxygenase inhibitors](#)
- 24 [6,875,784](#) [T](#) [Antimicrobial \[3.1.0.\] bicyclic oxazolidinone derivatives](#)
- 25 [6,872,725](#) [T](#) [Solid-state forms of N-\(2-hydroxyacetyl\)-5-\(4-piperidyl\)-4-\(4-pyrimidinyl\)-3-\(4-chlorophenyl\) pyrazole](#)
- 26 [6,870,070](#) [T](#) [Process to prepare 2-aminoindan derivatives](#)
- 27 [6,870,056](#) [T](#) [Substituted polycyclic aryl and heteroaryl pyridones useful for selective inhibition of the coagulation cascade](#)
- 28 [6,869,965](#) [T](#) [Antimicrobial quinolone derivatives and use of the same to treat bacterial infections](#)
- 29 [6,869,951](#) [T](#) [Method of changing conformation of a matrix metalloproteinase](#)
- 30 [6,867,217](#) [T](#) [Substituted polycyclic aryl and heteroaryl pyridones useful for selective inhibition of the coagulation cascade](#)
- 31 [6,867,018](#) [T](#) [Alzheimer's disease secretase, APP substrates therefor, and uses thereof](#)
- 32 [6,864,373](#) [T](#) [Stable amorphous celecoxib composite and process therefor](#)
- 33 [6,864,290](#) [T](#) [Statine derivatives for the treatment of Alzheimer's disease](#)
- 34 [6,864,080](#) [T](#) [Crystallization and structure of Staphylococcus aureus peptide deformylase](#)
- 35 [6,863,647](#) [T](#) [2-Ureido-thiazole derivatives, process for their preparation, and their use as antitumor agents](#)
- 36 [6,861,521](#) [T](#) [Stereoselective transacetalization of steroidal C-22 acetonide](#)
- 37 [6,861,433](#) [T](#) [Oxazolidinone photoaffinity probes](#)
- 38 [6,861,420](#) [T](#) [N-substituted 1,2,4,5-tetrahydro-1H-benzo\[d\]azepine compounds](#)
- 39 [6,858,846](#) [T](#) [Delivery device and method for its operation](#)
- 40 [6,858,635](#) [T](#) [Oxazolidinone photoaffinity probes](#)
- 41 [6,858,385](#) [T](#) [Pseudorabies virus protein](#)
- 42 [6,855,709](#) [T](#) [Pyridyl sulfone derivatives](#)
- 43 [6,852,761](#) [T](#) [Polycyclic aryl and heteroaryl substituted benzenes useful for selective inhibition of the coagulation cascade](#)
- 44 [6,852,753](#) [T](#) [Alkyl/aryl hydroxy or keto thiepine compounds as inhibitors of apical sodium co-dependent bile acid transport \(ASBT\) and taurocholate uptake](#)
- 45 [6,849,750](#) [T](#) [Processes and intermediates for preparing benzyl epoxides](#)
- 46 [6,849,669](#) [T](#) [Hydrophilic macromolecular compounds](#)
- 47 [6,849,653](#) [T](#) [Substituted pyrazolyl benzenesulfamide compounds for the treatment of inflammation](#)
- 48 [6,849,640](#) [T](#) [Therapeutic 1H-pyrido \[4,3-b\] indoles](#)
- 49 [6,849,616](#) [T](#) [Methods to potentiate intravenous estramustine phosphate](#)
- 50 [6,846,875](#) [T](#) [Hydrogels and methods for their production](#)
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Results of Search in 1976 to present db for:
(AN/pharmacia AND crystalline): 250 patents.
Hits 1 through 50 out of 250

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PAT. NO.	Title
1 6,903,097	T Heterocycle carboxamides as antiviral agents
2 6,900,244	T Anilino liver X-receptor modulators
3 6,897,318	T Process for making substituted pyrazoles
4 6,894,042	T Azabicyclic compounds for the treatment of disease
5 6,891,025	T Modified cytokine receptor protein
6 6,890,937	T Aromatic sulfone hydroxamic acid metalloprotease inhibitor
7 6,890,928	T Aromatic sulfone hydroxamic acids and their use as protease inhibitors
8 6,884,813	T Antimicrobial dihydrothiazine and dihydrothiopyran oxazolidinones
9 6,878,823	T 1,2,3,4,5,6-hexahydroazepino[4,5-b]indoles containing arylsulfones at the 9-position
10 6,872,725	T Solid-state forms of N-(2-hydroxyacetyl)-5-(4-piperidyl)-4-(4-pyrimidinyl)-3-(4-chlorophenyl) pyrazole
11 6,870,056	T Substituted polycyclic aryl and heteroaryl pyridones useful for selective inhibition of the coagulation cascade
12 6,869,965	T Antimicrobial quinolone derivatives and use of the same to treat bacterial infections
13 6,867,217	T Substituted polycyclic aryl and heteroaryl pyridones useful for selective inhibition of the coagulation cascade
14 6,864,373	T Stable amorphous celecoxib composite and process therefor
15 6,864,290	T Statine derivatives for the treatment of Alzheimer's disease
16 6,864,080	T Crystallization and structure of Staphylococcus aureus peptide deformylase
17 6,849,750	T Processes and intermediates for preparing benzyl epoxides
18 6,849,669	T Hydrophilic macromolecular compounds

- 19 [6,846,813](#) **T** [Compounds to treat alzheimer's disease](#)
- 20 [6,841,557](#) **T** [Compounds for the treatment of addictive disorders](#)
- 21 [6,833,453](#) **T** [Methods of producing oxazolidinone compounds](#)
- 22 [6,822,102](#) **T** [Dihydrobenzopyrans, dihydrobenzothiopyrans, and tetrahydroquinolines for the treatment of COX-2 mediated disorders](#)
- 23 [6,821,969](#) **T** [Thioxazinoquinolones as antiviral agents](#)
- 24 [6,815,460](#) **T** [Process for preparing prodrugs of benzenesulfonamide-containing cox-2 inhibitors](#)
- 25 [6,814,319](#) **T** [Laboratory scale milling process](#)
- 26 [6,800,646](#) **T** [Sulfamato hydroxamic acid metalloprotease inhibitor](#)
- 27 [6,800,635](#) **T** [Crystalline form II of cabergoline](#)
- 28 [6,797,003](#) **T** [Aspheric soft lens](#)
- 29 [6,796,975](#) **T** [Container for linezolid intravenous solution](#)
- 30 [6,794,544](#) **T** [Method for the preparation of tetrahydrobenzothiepinines](#)
- 31 [6,790,998](#) **T** [Phenyl inden-1-one compounds](#)
- 32 [6,767,934](#) **T** [Macromolecular compounds](#)
- 33 [6,765,003](#) **T** [3-Arylsulfonyl-2 \(substituted methyl\) propanoic acid derivatives as matrix metalloproteinase inhibitors](#)
- 34 [6,756,406](#) **T** [2-amino-2-alkyl-4 hexenoic and hexynoic acid derivatives useful as nitric oxide synthase inhibitors](#)
- 35 [6,750,342](#) **T** [Substituted polycyclic aryl and heteroaryl pyrimidinones useful for selective inhibition of the coagulation cascade](#)
- 36 [6,750,338](#) **T** [Process for making 5-substituted pyrazoles](#)
- 37 [6,750,233](#) **T** [Aromatic sulfone hydroxamic acid metalloprotease inhibitor](#)
- 38 [6,750,228](#) **T** [Aromatic sulfone hydroxamic acid metalloprotease inhibitor](#)
- 39 [6,747,090](#) **T** [Compositions capable of forming hydrogels in the eye](#)
- 40 [6,747,027](#) **T** [Thiol sulfonamide metalloprotease inhibitors](#)
- 41 [6,743,811](#) **T** [Oxazalidinone compounds and methods of preparation and use thereof](#)
- 42 [6,737,496](#) **T** [Photocurable siloxane polymers](#)
- 43 [6,727,363](#) **T** [Process for preparing crystalline form I of cabergoline](#)
- 44 [6,716,870](#) **T** [Prodrugs of 3-\(pyrrol-2-ylmethylidene\)-2-indolinone derivatives](#)
- 45 [6,716,829](#) **T** [Aldosterone antagonist and cyclooxygenase-2 inhibitor combination therapy to prevent or treat inflammation-related cardiovascular disorders](#)
- 46 [6,713,260](#) **T** [Methods of identifying compounds that bind to target species under isothermal denaturing conditions](#)
- 47 [6,710,067](#) **T** [Mannich base prodrugs of 3-\(pyrrol-2-ylmethylidene\)-2-indolinone derivatives](#)
- 48 [6,705,729](#) **T** [Methods of obtaining ophthalmic lenses providing the eye with reduced aberrations](#)
- 49 [6,696,449](#) **T** [Sulfonyl aryl hydroxamates and their use as matrix metalloprotease inhibitors](#)
- 50 [6,689,916](#) **T** [Phenyl propenone compounds](#)
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((AN/pharmacia AND crystalline) AND steroid): 23 patents.

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





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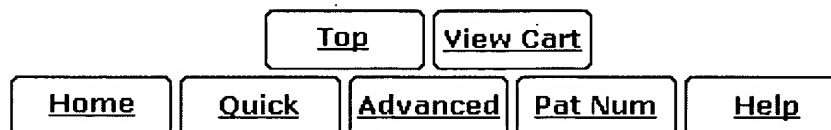
Refine Search

an/pharmacia and crystalline and steroid

PAT.
NO. Title

- 1 [6,878,823](#) [T](#) [1,2,3,4,5,6-hexahydroazepino\[4,5-b\]indoles containing arylsulfones at the 9-position](#)
- 2 [6,846,813](#) [T](#) [Compounds to treat alzheimer's disease](#)
- 3 [6,716,829](#) [T](#) [Aldosterone antagonist and cyclooxygenase-2 inhibitor combination therapy to prevent or treat inflammation-related cardiovascular disorders](#)
- 4 [6,653,308](#) [T](#) [3-\(4-amidopyrrol-2-ylmethylidene\)-2-indolinone derivatives as protein kinase inhibitors](#)
- 5 [6,613,880](#) [T](#) [Pipelicolic acid derivatives of proline threonine amides useful for the treatment of rheumatoid arthritis](#)
- 6 [6,573,293](#) [T](#) [Pyrrole substituted 2-indolinone protein kinase inhibitors](#)
- 7 [6,468,999](#) [T](#) [1,2,3,4,5,6,-hexahydroazepino \[4,5-b\]indoles containing arylsulfones at the 9-position](#)
- 8 [6,207,178](#) [T](#) [Solid lipid particles, particles of bioactive agents and methods for the manufacture and use thereof](#)
- 9 [6,143,853](#) [T](#) [Method for separation and synthetic polymers that can be used as separation media in the method](#)
- 10 [6,133,446](#) [T](#) [Heterocyclic compounds for the treatment of CNS and cardiovascular disorders](#)
- 11 [6,074,657](#) [T](#) [Administration of an injectable antibiotic in the ear of an animal](#)
- 12 [5,886,180](#) [T](#) [25-methylene and 24-25 -epoxy marcfortines and paraherquamides](#)
- 13 [5,877,317](#) [T](#) [Heterocyclic compounds for the treatment of CNS and cardiovascular disorders](#)
- 14 [5,795,986](#) [T](#) [Pyrimido>4,5-B!indoles](#)
- 15 [5,785,976](#) [T](#) [Solid lipid particles, particles of bioactive agents and methods for the manufacture and use thereof](#)
- 16 [5,759,404](#) [T](#) [Method for separation and synthetic polymers that can be used as separation media in the method](#)
- 17 [5,750,695](#) [T](#) [Antiparasitic paraherquamides](#)

- 18 [5,707,634](#)  [Finely divided solid crystalline powders via precipitation into an anti-solvent](#)
- 19 [5,703,078](#)  [Antiparasitic marcfortines and paraherquamides](#)
- 20 [5,686,610](#)  [Pyridyl piperazine compound](#)
- 21 [5,668,282](#)  [4,5-cyclicimidazo\[1,5-A\]quinoxalines](#)
- 22 [5,492,840](#)  [Surface plasmon resonance sensor unit and its use in biosensor systems](#)
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eplerenone

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MedlinePlus Drug Information: Eplerenone

Eplerenone is used alone or in combination with other medications to treat ... **Eplerenone** is in a class of medications called mineralocorticoid receptor ...

www.nlm.nih.gov/medlineplus/druginfo/medmaster/a603004.html - 23k - [Cached](#) - [Similar pages](#)

MedlinePlus Drug Information: Eplerenone (Systemic)

Pregnancy—**eplerenone** has not been studied in pregnant women. ... Breast-feeding—It is not known whether **eplerenone** passes into breast milk. ...

www.nlm.nih.gov/medlineplus/druginfo/uspdi/500431.html - 26k - [Cached](#) - [Similar pages](#)

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NEJM -- Eplerenone, a Selective Aldosterone Blocker, in Patients ...

Original Article from The New England Journal of Medicine -- **Eplerenone**, a Selective Aldosterone Blocker, in Patients with Left Ventricular Dysfunction ...

content.nejm.org/cgi/content/short/348/14/1309 - [Similar pages](#)

[PDF] 3 INSPRA eplerenone tablets DESCRIPTION

File Format: PDF/Adobe Acrobat - [View as HTML](#)

INSPRA contains **eplerenone**, a blocker of aldosterone binding at the mineralocorticoid ...

Eplerenone is an odorless, white to off-white crystalline powder. ...

www.fda.gov/cder/foi/label/2002/214371bl.pdf - [Similar pages](#)

eplerenone Consumer Drug Information

eplerenone Drug Information from Drugs.com. Includes side effects, interactions, indications.

www.drugs.com/MTM/eplerenone.html - 54k - [Cached](#) - [Similar pages](#)

Newly Approved Drug Therapies (805): Inspra (eplerenone tablets ...

Listing for Inspra (**eplerenone** tablets) in FDA Approved Drug Therapies Archives from CenterWatch Clinical Trials Listing Service.

www.centerwatch.com/patient/drugs/dru805.html - 17k - [Cached](#) - [Similar pages](#)

Eplerenone's indications expanded to include post-MI heart failure

Eplerenone's indications expanded to include post-MI heart failure. FDA has approved **eplerenone** (Inspra-Pfizer) for improving the survival of stable ...

www.pharmacist.com/articles/h_ts_0387.cfm - 12k - [Cached](#) - [Similar pages](#)

EHS: Mosby's Drug Consult - Drug Updates - Eplerenone 003570

Inspra contains **eplerenone**, a blocker of aldosterone binding at the ... Inspra for oral administration contains 25, 50, or 100 mg of **eplerenone** and the ...

www.mosbysdrugconsult.com/DrugConsult/003570.html - 66k - [Cached](#) - [Similar pages](#)

EPLERENONE - ORAL (Inspra) side effects, medical uses, and drug ...

Consumer information about the medication **EPLERENONE** - ORAL (Inspra), includes side effects, drug interactions, recommended dosages, and storage information ...

www.medicinenet.com/eplerenone-oral/article.htm - 37k - [Cached](#) - [Similar pages](#)

Yahoo! Health Drug Guide: Eplerenone Overview

Eplerenone may also be used for purposes other than those listed in this ... **Eplerenone** is available with a prescription under the brand name Inspra. ...

health.yahoo.com/drug/d04815a1 - 40k - [Cached](#) - [Similar pages](#)

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eplerenone

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PAT. NO.	Title
1 6,747,020	T Methods of treating heart failure and hypertension using combinations of eplerenone and an angiotensin converting enzyme inhibitor
2 6,716,829	T Aldosterone antagonist and cyclooxygenase-2 inhibitor combination therapy to prevent or treat inflammation-related cardiovascular disorders

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